10/019,249

file apry

ACCESSION NUMBER:

2000:209087 CAPLUS

DOCUMENT NUMBER:

132:343200

TITLE:

Parallel synthesis of a series of subtype-selective

NMDA receptor antagonists

AUTHOR(S):

Gregory, Tracy F.; Wright, Jon L.; Wise, Lawrence D.;

Meltzer, Leonard T.; Serpa, Kevin A.; Konkoy, Christopher S.; Whittemore, Edward R.; Woodward,

Richard M.

CORPORATE SOURCE:

Department of Chemistry, Division of Warner-Lambert Company, Parke-Davis Pharmaceutical Research, Ann

Arbor, MI, 48105, USA

SOURCE:

Bioorganic & Medicinal Chemistry Letters (2000),

10(6), 527-529

CODEN: BMCLE8; ISSN: 0960-894X

PUBLISHER:

Elsevier Science Ltd.

DOCUMENT TYPE:

Journal

LANGUAGE:

English

AΒ A series of 1-[(heteroarylthio)alkyl]-4-benzylpiperidines was rapidly synthesized through the use of parallel synthesis to investigate the binding affinity for the NR1A/2B receptor subtype.

IT 269079-52-9P 269079-54-1P

> RL: CAT (Catalyst use); SPN (Synthetic preparation); PREP (Preparation); USES (Uses)

(parallel synthesis and NR1A/2B receptor potency of

[(heteroarylthio)alkyl]benzylpiperidines as NMDA antagonists)

RN 269079-52-9 CAPLUS

CN 1H-Benzimidazole, 5-nitro-2-[[3-[4-(phenylmethyl)-1piperidinyl]propyl]thio]- (9CI) (CA INDEX NAME)

RN 269079-54-1 CAPLUS

CN 1H-Benzimidazole, 2-[[3-[4-(phenylmethyl)-1-piperidinyl]propyl]thio]-(9CI) (CA INDEX NAME)

REFERENCE COUNT:

3 THERE ARE 3 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT => file caplus FILE 'CAPLUS' ENTERED AT 13:56:58 ON 13 DEC 2002 USE IS SUBJECT TO THE TERMS OF YOUR STN CUSTOMER AGREEMENT. PLEASE SEE "HELP USAGETERMS" FOR DETAILS. COPYRIGHT (C) 2002 AMERICAN CHEMICAL SOCIETY (ACS)

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FILE COVERS 1907 - 13 Dec 2002 VOL 137 ISS 25 FILE LAST UPDATED: 12 Dec 2002 (20021212/ED)

This file contains CAS Registry Numbers for easy and accurate substance identification.

CAS roles have been modified effective December 16, 2001. Please check your SDI profiles to see if they need to be revised. For information on CAS roles, enter HELP ROLES at an arrow prompt or use the CAS Roles thesaurus (/RL field) in this file.

=> d que L1

STR

 $[CH_2]_{3-9}$

Structure attributes must be viewed using STN Express query preparation.

L3 33 SEA FILE=REGISTRY SSS FUL L1

L4 3 SEA FILE=CAPLUS L3

=> d 14 1-3 ibib abs hitstr

L4 ANSWER 1 OF 3 CAPLUS COPYRIGHT 2002 ACS ACCESSION NUMBER: 2001:12446 CAPLUS

DOCUMENT NUMBER: 134:86250

TITLE: Preparation and effect of benzimidazole compounds as

antiarteriosclerotics

INVENTOR(S): Aoki, Kozo; Aikawa, Kazuhiro; Kawakami, Masayuki

PATENT ASSIGNEE(S): Fuji Photo Film Co., Ltd., Japan

SOURCE: PCT Int. Appl., 37 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: Japanese

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

```
APPLICATION NO.
                                                                             DATE
      PATENT NO.
                            KIND
                                  DATE
                                                                            _____
                                                       _____
                           ____
                                   _____
                                                      WO 2000-JP4204
                                                                             20000627
      WO 2001000613
                            A1
                                   20010104
           W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN,
                CR, CU, CZ, DE, DK, DM, DZ, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NO, NZ, PL, PT, RO, RU, SD,
                SE, SG, SI, SK, SL, TJ, TM, TR, TT, TZ, UA, UG, US, UZ, VN, YU, ZA, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM
           RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZW, AT, BE, CH, CY,
               DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, BF, BJ, CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG
                             A1
                                   20020502
                                                     EP 2000-940847 20000627
               AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT,
                IE, SI, LT, LV, FI, RO, MK, CY, AL
PRIORITY APPLN. INFO.:
                                                   JP 1999-185568
                                                                         A 19990630
                                                   WO 2000-JP4204
                                                                         W 20000627
                               MARPAT 134:86250
OTHER SOURCE(S):
GΙ
```

$$\begin{array}{c|c}
 & O \\
 & R3 \\
 & X \\
 & R2 \\
 & X
\end{array}$$

AB Title compds.[I; wherein R1 is a substituent on the benzene ring which is selected from the group consisting of hydrogen, halogeno, lower alky, and lower alkoxy; R2 is hydrogen, alkyl, or acyl; and R3 is a substituent on the ring contg. nitrogen and Z; Z is a divalent group constituting a five-or six-membered ring; L is C4-C8 alkylene or an ethylene-oxy group represented by the general formula: (CH2CH2O)nCH2CH2 (wherein n is 1 or 2); and X is O or S] and salts thereof, which exhibit an inhibitory activity against the loading of macrophages in foam cells formation and are useful as the active ingredient of drugs to be used in the prevention and/or treatment of arteriosclerosis. Thus, the title compd. II was prepd. and tested.

II

IT 316362-98-8P

RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(prepn. and effect of benzimidazole compds. as antiarteriosclerotics)

RN 316362-98-8 CAPLUS

2,6-Piperidinedione, 1-[5-(1H-benzimidazol-2-ylthio)pentyl]- (9CI) (CA CN INDEX NAME)

REFERENCE COUNT:

13 THERE ARE 13 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

ANSWER 2 OF 3 CAPLUS COPYRIGHT 2002 ACS

ACCESSION NUMBER:

2001:12426 CAPLUS

DOCUMENT NUMBER:

134:86247

TITLE:

Preparation and effect of benzimidazoles as

antiarteriosclerotics

INVENTOR(S):

Aoki, Kozo; Aikawa, Kazuhiro; Kawakami, Masayuki; Yan,

Yongzhe

PATENT ASSIGNEE(S):

Fuji Photo Film Co., Ltd., Japan

SOURCE:

PCT Int. Appl., 52 pp. CODEN: PIXXD2

DOCUMENT TYPE:

Patent

LANGUAGE:

Japanese

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PA:	rent	NO.		KI	ND	DATE			A	PPLI	CATI	ои ис	ο.	DATE			
WO	2001	0005	88	A	1	2001	0104		W	20	00-J	P420	3	2000	0627		
	W:	ΑE,	AG,	AL,	AM,	AT,	AU,	AZ,	BA,	BB,	BG,	BR,	BY,	BZ,	CA,	CH,	CN,
		CR,	CU,	CZ,	DE,	DK,	DM,	DZ,	EE,	ES,	FI,	GB,	GD,	GE,	GH,	GM,	HR,
		HU,	ID,	IL,	IN,	IS,	JP,	KE,	KG,	KR,	KZ,	LC,	LK,	LR,	LS,	LT,	LU,
		LV,	MA,	MD,	MG,	MK,	MN,	MW,	MX,	MZ,	NO,	NZ,	PL,	PT,	RO,	RU,	SD,
		SE,	SG,	SI,	SK,	SL,	ТJ,	TM,	TR,	TT,	TZ,	UA,	UG,	US,	UZ,	VN,	YU,
		ZA,	ZW,	AM,	ΑZ,	BY,	KG,	KZ,	MD,	RU,	ТJ,	TM					
	RW:	GH,	GM,	ΚE,	LS,	MW,	MZ,	SD,	SL,	SZ,	TZ,	ŪĠ,	ZW,	ΑT,	BE,	CH,	CY,
		DE,	DK,	ES,	FI,	FR,	GB,	GR,	IE,	IT,	LU,	MC,	NL,	PT,	SE,	BF,	ВJ,
		CF,	CG,	CI,	CM,	GΑ,	GN,	GW,	ML,	MR,	NE,	SN,	TD,	TG			
EP	EP 1197487 A1 20020417						EP 2000-939171 20000627										
	R:	ΑT,	BE,	CH,	DE,	DK,	ES,	FR,	GB,	GR,	IT,	LI,	LU,	NL,	SE,	MC,	PT,
		ΙE,	SI,	LT,	LV,	FI,	RO										
RIT	RITY APPLN. INFO.:								JP 19	999-	1811	42	Α	1999	0628		
									30 0	200	TD 40	2.2	T.7	2000	2627		

PRIOR

WO 2000-JP4203 W 20000627

OTHER SOURCE(S):

MARPAT 134:86247

GI

AB Title compds. [I; wherein R1 is hydrogen, halogeno, lower alkyl, or lower alkoxy; R2 is hydrogen, alkyl, or acyl; R3 is a substituent on the ring or forming fused ring; A is O or CH2, or alternatively A represents a CH group binding to an adjacent carbon atom through a double bond; L is C4-C8 alkylene or an ethylene-oxy connecting group represented by the general formula: (CH2CH2O)nCH2CH2 (wherein n is 1 or 2); X is O, S, or methylene; and m is O or 1] or salts thereof, which exhibit inhibitory activities against the loading of macrophages in forming foam cells and are useful as the active ingredient of drugs to be used in the prevention and/or treatment of arteriosclerosis. Thus, the title compd. II was prepd. and tested.

IT 316371-85-4P 316371-87-6P 316371-89-8P 316371-92-3P 316371-94-5P 316371-97-8P 316371-98-9P 316372-05-1P 316372-06-2P 316372-07-3P 316372-10-8P 316372-11-9P 316372-12-0P 316372-13-1P 316372-14-2P 316372-15-3P 316372-22-2P 316372-23-3P 316372-24-4P 316372-25-5P 316372-26-6P 316372-27-7P 316372-28-8P 316372-29-9P 316372-30-2P 316372-31-3P 316372-32-4P 316372-52-8P

RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(prepn. and effect of benzimidazoles as antiarteriosclerotics) 316371-85-4 CAPLUS

1H-Benzimidazole, 2-[[5-(1-piperidinyl)pentyl]thio]- (9CI) (CA INDEX. NAME)

RN

CN

RN 316371-87-6 CAPLUS

CN 1H-Benzimidazole, 2-[[5-(4-methyl-1-piperidinyl)pentyl]thio]- (9CI) (CA INDEX NAME)

RN 316371-89-8 CAPLUS

CN 4-Piperidinecarboxylic acid, 1-[5-(1H-benzimidazol-2-ylthio)pentyl]-, ethyl ester (9CI) (CA INDEX NAME)

RN 316371-92-3 CAPLUS

CN 4-Piperidinol, 1-[5-(1H-benzimidazol-2-ylthio)pentyl]- (9CI) (CA INDEX NAME)

RN 316371-94-5 CAPLUS

CN 1H-Benzimidazole, 2-[[6-(4-methyl-1-piperidinyl)hexyl]thio]- (9CI) (CA INDEX NAME)

RN 316371-97-8 CAPLUS

CN 1H-Benzimidazole, 2-[(5-[1,4'-bipiperidin]-1'-ylpentyl)thio]- (9CI) (CA INDEX NAME)

RN 316371-98-9 CAPLUS

CN 1H-Benzimidazole, 2-[[5-(3,5-dimethyl-1-piperidinyl)pentyl]thio]- (9CI) (CA INDEX NAME)

RN 316371-99-0 CAPLUS

CN 4-Piperidineethanol, 1-[5-(1H-benzimidazol-2-ylthio)pentyl]- (9CI) (CA INDEX NAME)

RN 316372-05-1 CAPLUS

CN 1H-Benzimidazole, 2-[[5-(2-methyl-1-piperidinyl)pentyl]thio]- (9CI) (CA INDEX NAME)

RN 316372-06-2 CAPLUS

CN 3-Piperidinecarboxamide, 1-[5-(1H-benzimidazol-2-ylthio)pentyl]- (9CI) (CA INDEX NAME)

RN 316372-07-3 CAPLUS

CN 1H-Benzimidazole, 2-[[4-(4-methyl-1-piperidinyl)butyl]thio]- (9CI) (CA INDEX NAME)

RN 316372-10-8 CAPLUS

CN 1H-Benzimidazole, 2-[[5-(2,6-dimethyl-1-piperidinyl)pentyl]thio]- (9CI) (CA INDEX NAME)

$$N$$
 $S-(CH_2)_5-N$
 Me
 Me
 Me

RN 316372-11-9 CAPLUS

CN 3-Piperidinemethanol, 1-[5-(1H-benzimidazol-2-ylthio)pentyl]- (9CI) (CA INDEX NAME)

RN 316372-12-0 CAPLUS

CN 2-Piperidinemethanol, 1-[5-(1H-benzimidazol-2-ylthio)pentyl]- (9CI) (CA INDEX NAME)

RN 316372-13-1 CAPLUS

CN 2-Piperidineethanol, 1-[5-(1H-benzimidazol-2-ylthio)pentyl]- (9CI) (CA INDEX NAME)

RN 316372-14-2 CAPLUS

CN 3-Piperidinol, 1-[5-(1H-benzimidazol-2-ylthio)pentyl]- (9CI) (CA INDEX NAME)

RN 316372-15-3 CAPLUS

CN 1H-Benzimidazole, 2-[[8-(4-methyl-1-piperidinyl)octyl]thio]- (9CI) (CA INDEX NAME)

RN 316372-22-2 CAPLUS

CN 4-Piperidinecarbonitrile, 1-[5-(1H-benzimidazol-2-ylthio)pentyl]-4-phenyl-(9CI) (CA INDEX NAME)

RN 316372-23-3 CAPLUS

CN 4-Piperidinecarbonitrile, 4-acetyl-1-[5-(1H-benzimidazol-2-ylthio)pentyl]-(9CI) (CA INDEX NAME)

RN 316372-24-4 CAPLUS

CN 4-Piperidinecarbonitrile, 1-[5-(1H-benzimidazol-2-ylthio)pentyl]-4-hydroxy-(9CI) (CA INDEX NAME)

RN 316372-25-5 CAPLUS

CN 4-Piperidinecarbonitrile, 1-[4-(1H-benzimidazol-2-ylthio)butyl]-4-phenyl-(9CI) (CA INDEX NAME)

$$N$$
 S- (CH₂)₄ N N CN

RN 316372-26-6 CAPLUS

CN 4-Piperidinecarbonitrile, 1-[6-(1H-benzimidazol-2-ylthio)hexyl]-4-phenyl-(9CI) (CA INDEX NAME)

RN 316372-27-7 CAPLUS

CN 4-Piperidinecarbonitrile, 1-[8-(1H-benzimidazol-2-ylthio)octyl]-4-phenyl-(9CI) (CA INDEX NAME)

RN 316372-28-8 CAPLUS

CN 4-Piperidinol, 1-[5-(1H-benzimidazol-2-ylthio)pentyl]-4-(4-chlorophenyl)-(9CI) (CA INDEX NAME)

RN 316372-29-9 CAPLUS

CN 4-Piperidinol, 1-[6-(1H-benzimidazol-2-ylthio)hexyl]-4-phenyl- (9CI) (CA INDEX NAME)

RN 316372-30-2 CAPLUS

CN 4-Piperidinol, 1-[4-(1H-benzimidazol-2-ylthio)butyl]-4-phenyl- (9CI) (CA INDEX NAME)

RN 316372-31-3 CAPLUS

CN 4-Piperidinol, 1-[5-(1H-benzimidazol-2-ylthio)pentyl]-4-(phenylmethyl)-(9CI) (CA INDEX NAME)

RN 316372-32-4 CAPLUS

CN 1H-Benzimidazole, 2-[[5-(4-methoxy-4-phenyl-1-piperidinyl)pentyl]thio]-(9CI) (CA INDEX NAME)

316372-52-8 CAPLUS RN

1H-Benzimidazole, 2-[[5-(3-methyl-1-piperidinyl)pentyl]thio]- (9CI) CN INDEX NAME)

REFERENCE COUNT:

2 THERE ARE 2 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

ANSWER 3 OF 3 CAPLUS COPYRIGHT 2002 ACS

ACCESSION NUMBER:

2000:209087 CAPLUS

DOCUMENT NUMBER:

132:343200

TITLE:

Parallel synthesis of a series of subtype-selective

NMDA receptor antagonists

AUTHOR(S):

Gregory, Tracy F.; Wright, Jon L.; Wise, Lawrence D.;

Meltzer, Leonard T.; Serpa, Kevin A.; Konkoy, Christopher S.; Whittemore, Edward R.; Woodward,

Richard M.

CORPORATE SOURCE:

Department of Chemistry, Division of Warner-Lambert

Company, Parke-Davis Pharmaceutical Research, Ann

Arbor, MI, 48105, USA

SOURCE:

Bioorganic & Medicinal Chemistry Letters (2000),

10(6), 527-529

CODEN: BMCLE8; ISSN: 0960-894X

PUBLISHER:

Elsevier Science Ltd. Journal

DOCUMENT TYPE:

English LANGUAGE:

AΒ A series of 1-[(heteroarylthio)alkyl]-4-benzylpiperidines was rapidly synthesized through the use of parallel synthesis to investigate the binding affinity for the NR1A/2B receptor subtype.

IT 269079-52-9P 269079-54-1P

> RL: CAT (Catalyst use); SPN (Synthetic preparation); PREP (Preparation); USES (Uses)

(parallel synthesis and NR1A/2B receptor potency of

[(heteroarylthio)alkyl]benzylpiperidines as NMDA antagonists)

RN 269079-52-9 CAPLUS

CN 1H-Benzimidazole, 5-nitro-2-[[3-[4-(phenylmethyl)-1piperidinyl]propyl]thio]- (9CI) (CA INDEX NAME)

RN 269079-54-1 CAPLUS

1H-Benzimidazole, 2-[[3-[4-(phenylmethyl)-1-piperidinyl]propyl]thio]-CN (9CI) (CA INDEX NAME)

REFERENCE COUNT:

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3

FILE 'USPAT2' ENTERED AT .13:58:03 ON 13 DEC 2002 CA INDEXING COPYRIGHT (C) 2002 AMERICAN CHEMICAL SOCIETY (ACS)

=> d que

L1 STR

$$\begin{bmatrix} CH_2 \end{bmatrix}_{3-9}$$

Structure attributes must be viewed using STN Express query preparation.

L3 33 SEA FILE=REGISTRY SSS FUL L1

L5 0 SEA L3

=> file caold
FILE 'CAOLD' ENTERED AT 13:58:13 ON 13 DEC 2002
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FILE COVERS 1907-1966

FILE LAST UPDATED: 01 May 1997 (19970501/UP)

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This file supports REG1stRY for direct browsing and searching of all substance data from the REGISTRY file. Enter HELP FIRST for more information.

=> s 13

L6 0 L3

=> file caplus

FILE 'CAPLUS' ENTERED AT 14:22:37 ON 13 DEC 2002

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FILE COVERS 1907 - 13 Dec 2002 VOL 137 ISS 25 FILE LAST UPDATED: 12 Dec 2002 (20021212/ED)

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CAS roles have been modified effective December 16, 2001. Please check your SDI profiles to see if they need to be revised. For information on CAS roles, enter HELP ROLES at an arrow prompt or use the CAS Roles thesaurus (/RL field) in this file.

=> d que

L1

STR

Structure attributes must be viewed using STN Express query preparation.

L3 1 SEA FILE=REGISTRY SSS FUL L1

L4 1 SEA FILE=CAPLUS L3

=> d 14 ibib abs hitstr

L4 ANSWER 1 OF 1 CAPLUS COPYRIGHT 2002 ACS ACCESSION NUMBER: 2001:12426 CAPLUS

DOCUMENT NUMBER: 134:86247

TITLE: Preparation and effect of benzimidazoles as

antiarteriosclerotics

INVENTOR(S): Aoki, Kozo; Aikawa, Kazuhiro; Kawakami, Masayuki; Yan,

Yongzhe

PATENT ASSIGNEE(S): Fuji Photo Film Co., Ltd., Japan

SOURCE: PCT Int. Appl., 52 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: Japanese

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

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APPLICATION NO.
     PATENT NO.
                     KIND DATE
    WO 2001000588
                           20010104
                                          WO 2000-JP4203
                                                           20000627
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            CR, CU, CZ, DE, DK, DM, DZ, EE, ES, FI, GB, GD, GE, GH, GM, HR,
            HU, ID, IL, IN, IS, JP, KE, KG, KR, KZ, LC, LK, LR, LS, LT, LU,
            LV, MA, MD, MG, MK, MN, MW, MX, MZ, NO, NZ, PL, PT, RO, RU, SD,
            SE, SG, SI, SK, SL, TJ, TM, TR, TT, TZ, UA, UG, US, UZ, VN, YU,
             ZA, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM
         RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZW, AT, BE, CH, CY,
            DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, BF, BJ,
            CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG
                      A1
                          20020417
                                         EP 2000-939171 20000627
     EP 1197487
         R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT,
             IE, SI, LT, LV, FI, RO
PRIORITY APPLN. INFO.:
                                        JP 1999-181142
                                                        A 19990628
                                                        W 20000627
                                       WO 2000-JP4203
OTHER SOURCE(S):
                       MARPAT 134:86247
GI
```

AB Title compds. [I; wherein R1 is hydrogen, halogeno, lower alkyl, or lower alkoxy; R2 is hydrogen, alkyl, or acyl; R3 is a substituent on the ring or forming fused ring; A is O or CH2, or alternatively A represents a CH group binding to an adjacent carbon atom through a double bond; L is C4-C8 alkylene or an ethylene-oxy connecting group represented by the general formula: (CH2CH2O)nCH2CH2 (wherein n is 1 or 2); X is O, S, or methylene; and m is O or 1] or salts thereof, which exhibit inhibitory activities against the loading of macrophages in forming foam cells and are useful as the active ingredient of drugs to be used in the prevention and/or treatment of arteriosclerosis. Thus, the title compd. II was prepd. and tested.

Ι

IT 316371-96-7P

RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); SPN (Synthetic preparation); THU (Therapeutic use);

REFERENCE COUNT:

2 THERE ARE 2 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

=> file uspatall FILE 'USPATFULL' ENTERED AT 14:23:17 ON 13 DEC 2002 CA INDEXING COPYRIGHT (C) 2002 AMERICAN CHEMICAL SOCIETY (ACS)

FILE 'USPAT2' ENTERED AT 14:23:17 ON 13 DEC 2002 CA INDEXING COPYRIGHT (C) 2002 AMERICAN CHEMICAL SOCIETY (ACS)

=> s 13 L5 0 L3

=> file reg
FILE 'REGISTRY' ENTERED AT 14:25:14 ON 13 DEC 2002
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Property values tagged with IC are from the ZIC/VINITI data file provided by InfoChem.

STRUCTURE FILE UPDATES: 12 DEC 2002 HIGHEST RN 476148-76-2 DICTIONARY FILE UPDATES: 12 DEC 2002 HIGHEST RN 476148-76-2

TSCA INFORMATION NOW CURRENT THROUGH MAY 20, 2002

Please note that search-term pricing does apply when conducting SmartSELECT searches.

Crossover limits have been increased. See HELP CROSSOVER for details.

Experimental and calculated property data are now available. See HELP PROPERTIES for more information. See STNote 27, Searching Properties in the CAS Registry File, for complete details: http://www.cas.org/ONLINE/STN/STNOTES/stnotes27.pdf

=> d que L6 STR

G1 0, CH2

Structure attributes must be viewed using STN Express query preparation. L8 0 SEA FILE=REGISTRY SSS FUL L6

=> file caplus
FILE 'CAPLUS' ENTERED AT 14:26:48 ON 13 DEC 2002
USE IS SUBJECT TO THE TERMS OF YOUR STN CUSTOMER AGREEMENT.
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FILE COVERS 1907 - 13 Dec 2002 VOL 137 ISS 25 FILE LAST UPDATED: 12 Dec 2002 (20021212/ED)

This file contains CAS Registry Numbers for easy and accurate substance identification.

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G1 0, CH2

Structure attributes must be viewed using STN Express guery preparation.

10/019,249

L11 8 SEA FILE=REGISTRY SSS FUL L9

=> d l11 1-8 ibib abs hitstr => d que STR L9

G1 0,CH2

Structure attributes must be viewed using STN Express query preparation.

8 SEA FILE=REGISTRY SSS FUL L9 L11

L12 3 SEA FILE=CAPLUS L11

=> d 112 1-8 ibib abs hitstr

L12 ANSWER 1 OF 3 CAPLUS COPYRIGHT 2002 ACS

ACCESSION NUMBER:

2001:12426 CAPLUS

DOCUMENT NUMBER:

134:86247

TITLE:

Preparation and effect of benzimidazoles as

antiarteriosclerotics

INVENTOR(S):

Aoki, Kozo; Aikawa, Kazuhiro; Kawakami, Masayuki; Yan,

Yongzhe

PATENT ASSIGNEE(S):

Fuji Photo Film Co., Ltd., Japan

SOURCE:

PCT Int. Appl., 52 pp. CODEN: PIXXD2

DOCUMENT TYPE:

Patent

LANGUAGE:

Japanese

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

	PAT	ENT	NO.		KI	ND.	DATE			A	PPLI	CATI	ои ис	ο.	DATE			
	wo	2001	0005	88	A:	 1	2001	0104		W	20	 00-J1	P420:	3	20000	0627		
		W:													BZ,			
			CR,	CU,	CZ,	DE,	DK,	DM,	DZ,	EE,	ES,	FI,	GB,	GD,	GE,	GH,	GM,	HR,
			HU,	ID,	IL,	IN,	IS,	JP,	KE,	KG,	KR,	ΚZ,	LC,	LK,	LR,	LS,	LT,	LU,
			LV,	MA,	MD,	MG,	MK,	MN,	MW,	MX,	MZ,	NO,	NZ,	PL,	PT,	RO,	RU,	SD,
			SE,	SG,	SI,	SK,	SL,	ТJ,	TM,	TR,	TT,	TZ,	UA,	UG,	US,	UZ,	VN,	YU,
			ZA,	ZW,	AM,	ΑZ,	BY,	KG,	ΚZ,	MD,	RU,	ТJ,	TM					
		RW:	GH,	GM,	KE,	LS,	MW,	MZ,	SD,	SL,	SZ,	TZ,	UG,	ZW,	AT,	BE,	CH,	CY,
															PT,			
			CF,	CG,	CI,	CM,	GA,	GN,	GW,	ML,	MR,	NE,	SN,	TD,	TG			
	ΕP	1197	487		A.	1	2002	0417		E	P 20	00-93	3917:	1	20000	0627		
		R:	AT,	BE,	CH,	DE,	DK,	ES,	FR,	GB,	GR,	IT,	LI,	LU,	NL,	SE,	MC,	PT,
			ΙE,	SI,	LT,	LV,	FI,	RO										
PRIOR	ITY	APP	LN.	INFO	. :					JP 19	999-:	1811	42	Α	19990	0628		
									1	WO 20	000-	JP420	03	W	20000	0627		
OTHER	SC	URCE	(S):			MAR	PAT	134:8	3624	7								

GI

AB Title compds. [I; wherein Rl is hydrogen, halogeno, lower alkyl, or lower alkoxy; R2 is hydrogen, alkyl, or acyl; R3 is a substituent on the ring or forming fused ring; A is O or CH2, or alternatively A represents a CH group binding to an adjacent carbon atom through a double bond; L is C4-C8 alkylene or an ethylene-oxy connecting group represented by the general formula: (CH2CH2O)nCH2CH2 (wherein n is 1 or 2); X is O, S, or methylene; and m is O or 1] or salts thereof, which exhibit inhibitory activities against the loading of macrophages in forming foam cells and are useful as the active ingredient of drugs to be used in the prevention and/or treatment of arteriosclerosis. Thus, the title compd. II was prepd. and tested.

IT 316372-38-0P 316372-39-1P 316372-40-4P 316372-41-5P

RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(prepn. and effect of benzimidazoles as antiarteriosclerotics)

RN 316372-38-0 CAPLUS

CN 4-Piperidinecarbonitrile, 1-[5-(1H-benzimidazol-2-yloxy)pentyl]-4-phenyl-(9CI) (CA INDEX NAME)

RN 316372-39-1 CAPLUS

CN 4-Piperidinol, 1-[5-(1H-benzimidazol-2-yloxy)pentyl]-4-phenyl- (9CI) (CA INDEX NAME)

RN 316372-40-4 CAPLUS

CN 4-Piperidinecarbonitrile, 1-[6-(1H-benzimidazol-2-yl)hexyl]-4-phenyl-(9CI) (CA INDEX NAME)

RN 316372-41-5 CAPLUS

CN 4-Piperidinol, 1-[6-(1H-benzimidazol-2-yl)hexyl]-4-phenyl- (9CI) (CA INDEX NAME)

REFERENCE COUNT:

THERE ARE 2 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L12 ANSWER 2 OF 3 CAPLUS COPYRIGHT 2002 ACS

2

ACCESSION NUMBER:

1990:423912 CAPLUS

DOCUMENT NUMBER:

113:23912

TITLE:

Preparation of benzimidazole derivatives as

antihistaminics

INVENTOR(S):
PATENT ASSIGNEE(S):

Giani, Roberto; Parini, Ettore; Tonon, Giancarlo

Dompe Farmaceutici S.p.A., Italy

SOURCE:

Eur. Pat. Appl., 8 pp. CODEN: EPXXDW

DOCUMENT TYPE:

Patent

LANGUAGE:

English

FAMILY ACC. NUM. COUNT:

1

PATENT INFORMATION:

PAT	TENT NO.		KIND	DATE	APPLICATION NO. DATE
	350467 350467		A1 B1	19900110 19931229	EP 1989-830312 19890706
	R: AT,	BE,	CH, DE	, ES, FR,	
	02067272 05067624		A2 B4	19900307 19930927	JP 1989-173174 19890706
US	4971980		Α	19901120	US 1989-376075 19890706
ΑT	99295		E	19940115	AT 1989-830312 19890706
ES	2062098		Т3	19941216	ES 1989-830312 19890706

PRIORITY APPLN. INFO.:

IT 1988-21271 EP 1989-830312 19880707 19890706

OTHER SOURCE(S):

MARPAT 113:23912

Ι

GΙ

AB Title compds. I [A = CH2CHMe, CHMeCH2; X = PhCH2, fluorobenzyl, EtOCH2CH2, H2C: CHCH2OCH2CH2, tetrahydrofurfuryl; R1, R2 = (un) satd. C1-4 alkyl, R1R2N = (substituted) pyrrolidinyl-piperidinyl; m = 0-5; n = 0,1] and their salts, are prepd. 2-[3-(Dimethylamino)propyl]benzimidazole (prepn. given) in DMF was treated with NaH and warmed to 60.degree. to which EtOCH2CH2Cl was added and maintained at 60.degree. for 5 h to give 1-ethoxy-2-[3-(dimethylamino)propyl]benzimidazole (II). II had ED50 of 4.54 .mu.g/kg orally in guinea pigs against histamine-induced mortality.

IT 127841-94-5P

> RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(prepn. and reaction of, in prepn. of antihistaminics)

RN 127841-94-5 CAPLUS

1H-Benzimidazole, 2-[4-(1-piperidinyl)butyl]- (9CI) (CA INDEX NAME) CN

TT 127842-02-8P

> RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(prepn. of, as antihistaminic)

RN 127842-02-8 CAPLUS

1H-Benzimidazole, 1-(2-ethoxyethyl)-2-[4-(1-piperidinyl)butyl]- (9CI) CN (CA INDEX NAME)

L12 ANSWER 3 OF 3 CAPLUS COPYRIGHT 2002 ACS ACCESSION NUMBER:

1975:156308 CAPLUS

DOCUMENT NUMBER:

82:156308

TITLE:

Benzimidazole derivatives

INVENTOR(S):

Hasegawa, Hajime; Tsuda, Nobutada; Hasoya, Masahiro

PATENT ASSIGNEE(S):

Yoshitomi Pharmaceutical Industries, Ltd.

SOURCE:

Japan., 4 pp. CODEN: JAXXAD

DOCUMENT TYPE:

Patent

LANGUAGE:

Japanese

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATENT NO. KIND DATE APPLICATION NO. DATE ______ JP 49041198 19741107 JP 1970-26357 19700328 В4

GΙ For diagram(s), see printed CA Issue.

AΒ Twenty-three benzimidazoles [I, R = CH2Ph, CH2C6H4Cl-p, etc., R1 = O(CH2)3NMe2, OCH2CH2NMe2, 3-morpholinopropoxy, SCH2CH2NMe2, S(CH2)3NMe2, S(CH2)2NHMe, SCH2CH2N(CH2Ph)2, etc., R2 = H, 6-Cl, 5-MeO, etc.] or their salts, useful as antihistaminics, analgesics, and inflammation inhibitors (no data), were prepd. by treating the chloro deriv. (I, R1 = C1) with the appropriate alc. or thiol in the presence of NaH. For example, NaOCH2CH2NMe2 (obtained from 8.9 g HOCH2CH2NMe2 and 4.8 g NaH) was refluxed with I (R = CH2Ph, R1 = $\overline{\text{Cl}}$, R2 = H) (21.2 g) in benzene for 4 hr and the product treated with (CO2H)2 to give 20 g I (R = CH2Ph, R1 = OCH2CH2NMe2, R2 = H).cntdot.(CO2H)2.

IT 55415-22-0P

> RL: SPN (Synthetic preparation); PREP (Preparation) (prepn. of)

RN 55415-22-0 CAPLUS

CN 1H-Benzimidazole, 1-[(4-chlorophenyl)methyl]-5-methoxy-2-[3-(2-methyl-1piperidinyl)propoxy]-, ethanedioate (1:1) (9CI) (CA INDEX NAME)

CM 1

CRN 55415-21-9 CMF C24 H30 Cl N3 O2

CM 2

CRN 144-62-7 CMF C2 H2 O4

=> file uspatall

FILE 'USPATFULL' ENTERED AT 14:28:25 ON 13 DEC 2002

CA INDEXING COPYRIGHT (C) 2002 AMERICAN CHEMICAL SOCIETY (ACS)

FILE 'USPAT2' ENTERED AT 14:28:25 ON 13 DEC 2002

CA INDEXING COPYRIGHT (C) 2002 AMERICAN CHEMICAL SOCIETY (ACS)

=> d que

STR L9

$$\begin{bmatrix} CH_2 \end{bmatrix}_{3-9}$$

G1. O, CH2

Structure attributes must be viewed using STN Express query preparation.

8 SEA FILE=REGISTRY SSS FUL L9 L11

T.13 1 SEA L11

=> d 113 ibib abs hitstr

L13 ANSWER 1 OF 1 USPATFULL

ACCESSION NUMBER:

90:89303 USPATFULL

TITLE:

Pharmacologically active benzimidazole derivatives

INVENTOR(S):

Roberto, Giani P., Milan, Italy Ettore, Parini, Milan, Italy Giancarlo, Tonon, Milan, Italy

PATENT ASSIGNEE(S):

Dompe Farmaceutici S.p.A., Milan, Italy (non-U.S.

corporation)

NUMBER KIND DATE

PATENT INFORMATION:

US 4971980 19901120

APPLICATION INFO.:

US 1989-376075 19890706 (7)

DATE NUMBER -----

PRIORITY INFORMATION:

IT 1988-21271 19880707

DOCUMENT TYPE:

Utility

FILE SEGMENT:

Granted

PRIMARY EXAMINER:

Lee: Mary C.

ASSISTANT EXAMINER:

LEGAL REPRESENTATIVE:

Haley, Jacqueline

Armstrong, Nikaido, Marmelstein, Kubovcik & Murray

NUMBER OF CLAIMS:

EXEMPLARY CLAIM:

1,6

LINE COUNT:

312

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

Novel benzimidazole derivatives (I) are described, of formula ##STR1## AB wherein A represents ##STR2## n is 0 or 1; m represents 0 or an integer of from 1 to 5 inclusive, provided that when n is 0, m represents an integer of from 2 to 5 inclusive;

X represents a radical selected from the group consisting of benzyl, fluorobenzyl, alkoxyalkyl and tetrahydrofurfuryl;

R.sub.1 and R.sub.2 represent each a saturated or unsaturated alkyl radical having of from 1 to 4 carbon atoms or they may form, together with the adjacent nitrogen atom, an optionally substituted heterocyclic ring selected from the group consisting of pyrrolidine and piperidine, and the corresponding, non-toxic, pharmaceutically acceptable acid addition salts.

The compounds (I) are endowed with an interesting antihistaminic activity.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

IT 127841-94-5P

(prepn. and reaction of, in prepn. of antihistaminics)

RN 127841-94-5 USPATFULL

CN 1H-Benzimidazole, 2-[4-(1-piperidinyl)butyl]- (9CI) (CA INDEX NAME)

$$\begin{array}{c}
H \\
N \\
\end{array}$$
(CH₂) 4 — N

IT 127842-02-8P

(prepn. of, as antihistaminic)

RN 127842-02-8 USPATFULL

CN 1H-Benzimidazole, 1-(2-ethoxyethyl)-2-[4-(1-piperidinyl)butyl]- (9CI) (CA INDEX NAME)